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(54) Title: NOVEL USES FOR GAL4-RECEPTOR CON	STRU	CTS
(57) Abstract		

In accordance with the present invention, it has been discovered that chimeric receptors comprising at least the ligand binding domain of a member of the steroid/thyroid receptor protein superfamily having incorporated therein the DNA binding domain of the yeast GALA protein are capable of forming functional homodimers. Such homodimers are useful, for example, in assays to determine the ligand for an orphan member of the steroid/thyroid superfamily of receptors, for identifying compounds which are selective for a specific steroid/thyroid receptor protein from among a plurality of such receptors which respond to a common ligand, and the like.

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Novel Uses for GAL4-Receptor Constructs

ACKNOWLEDGEMENT

This invention was made with Government support under Grant Nos. HD27183, awarded by the National Institute of Health. The Government has certain rights in the invention.

FIELD OF THE INVENTION

The present invention relates to intracellular receptors, and ligands therefor. In a particular aspect, the present invention relates to methods for the identification of compounds which function as ligands (or ligand precursors) for intracellular receptors. In another aspect, the present invention relates to methods for the identification of compounds which are selective for a specific receptor protein from among a plurality of receptors which respond to a common ligand. In a further aspect, the present invention relates to novel chimeric intracellular receptors and uses therefor.

BACKGROUND OF THE INVENTION

A central problem in eukaryotic molecular biology continues to be the elucidation of molecules and mechanisms that mediate specific gene regulation. As part of the scientific attack on this problem, a great deal of work has been done in efforts to identify ligands (i.e., exogenous inducers) which are capable of mediating specific gene regulation.

Although much remains to be learned about the specifics of gene regulation, it is known that ligands modulate gene transcription by acting in concert with intracellular components, including intracellular receptors and discrete DNA sequences known as hormone response elements (HREs).

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As additional members of the steroid/thyroid superfamily of receptors are identified, the search for exogenous inducers for such newly discovered receptors (i.e., naturally occurring (or synthetic) inducers) has become an important part of the effort to learn about the specifics of gene regulation.

The identification of compounds which directly or indirectly interact with intracellular receptors, and thereby affect transcription of hormone-responsive genes, would be of significant value, e.g., for therapeutic applications.

Additional novel intracellular receptors (i.e., members of the steroid/thyroid superfamily of receptors) continue to be identified. Frequently, however, the primary ligand(s) for these novel receptors can not readily be identified. Accordingly, methods for the ready identification of ligands for such receptors would be of great value.

Of the intracellular receptors for which ligands 20 have been identified, there are some which respond to a common ligand (e.g., retinoic acid receptor (RAR) and retinoid X receptor (RXR) both respond to the presence of retinoic acid; glucocorticoid receptor (GR) and mineralocorticoid receptor (MR) both respond to the 25 presence of glucocorticoids, etc.). In order to modulate process(es) mediated by one, but not all of the receptor subtypes which respond to a common ligand, a great deal of energy has been invested in efforts to discover ligands which are selective for a single subtype. It is difficult, 30 however, to differentiate between the actions of a single ligand on several different receptor Accordingly, methods to identify compounds selective for a single subtype would also be of great value.

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It has recently been discovered that some intracellular receptors are functional for regulation of transcription only when associated with another member of the steroid/thyroid superfamily of receptors (i.e., as a 5 heterodimer). Until the physiological processes impacted by a specific receptor are known, however, it is not possible to determine if an orphan receptor functions as a homodimer or requires a partner (so as to form a functional Accordingly, heterodimer). means to simplify 10 characterization of orphan receptors (i.e., receptors for which no ligand has yet been identified), without the need to determine if said orphan requires a partner to produce a functional receptor, would also be of great value.

Other information helpful in the understanding 15 and practice of the present invention can be found in commonly assigned United States Patent Nos. 5,071,773 and 4,981,784; and United States Patent Application Nos. 325,240, filed March 17, 1989; 370,407, filed June 22, 1989; and 438,757, filed November 16, 1989, all of which 20 are hereby incorporated herein by reference in their entirety.

BRIEF DESCRIPTION OF THE INVENTION

In accordance with the present invention, we have discovered that chimeric receptors comprising a member of 25 the steroid/thyroid receptor protein superfamily having incorporated therein the DNA binding domain of the yeast GAL4 protein are capable of forming functional homodimers. Such homodimers are useful, for example, in assays to determine the ligand for an orphan member of the 30 steroid/thyroid superfamily of receptors, for identifying compounds which are selective for a steroid/thyroid receptor protein from among a plurality of such receptors which respond to a common ligand, and the like.

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Such assays have a broad dynamic range and very low background signals, since the response system triggered by the presence of suitable ligand is not endogenous to the cells employed for the assays. A response is obtained only when the chimeric receptors of the invention are activa d, without competition from endogenous receptors which may be present in the test cells.

In accordance with another embodiment of the invention, present are provided methods the identification of compounds which are convertible into ligand or ligand precursors for intracellular receptors, or other compounds which may induce receptors through a nonligand activated pathway. In accordance with yet another embodiment of the present invention, methods are provided 15 for the identification of compounds (e.g., enzymes) which are capable of converting appropriate precursor compounds ligands for intracellular receptors (or other compounds which may induce receptors through a non-ligand activated pathway). In accordance with still another 20 embodiment of the present invention, methods are provided for the identification of cell lines (or active fractions thereof) which are capable of promoting the above-described conversions.

BRIEF DESCRIPTION OF THE FIGURES

Figure 1 is a restriction map of the GAL4 encoding portion of vector pCMX-GAL4 employed in the preparation of GAL4-receptor fusion proteins.

Figure 2 is a restriction map of the linker set forth in SEQ ID NO:4.

Figure 3 is a dose response curve for various hRARα constructs, in the presence of all trans retinoic acid. In the figure, ◊ designates unmodified hRARα, ♦

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designates the chimeric receptor GRR (i.e., a chimeric receptor wherein the amino terminal domain is derived from the glucocorticoid receptor and the DNA and ligand binding domains are derived from RAR),

designates the chimeric receptor wherein the amino terminal domain and the DNA binding domain are derived from GR, and the ligand binding domain is derived from RAR), and designates GAL4-hRARa.

DETAILED DESCRIPTION OF THE INVENTION

In accordance with the present invention, there is provided a method to determine the ligand or ligand precursor for an orphan member of the steroid/thyroid receptor protein superfamily, said method comprising

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contacting cells containing a modified form of said orphan member with a putative ligand for said orphan member;

wherein said modified form of said orphan member is produced by introducing the DNA binding domain of GAL4 into said orphan member, and

wherein said cells contain a GAL4 response element operatively linked to a reporter gene, and thereafter

monitoring the expression of reporter gene product.

Optionally, the modified receptor employed in the practice of the present invention will also contain one or more exogenous transactivation domains, such as, for example, the τ_1 or τ_2 transactivation domains described in United States Patent No. 5,217,867, which is incorporated by reference herein in its entirety.

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Invention assay method is especially useful where the response element for said orphan receptor has not been identified and/or where the ability of said receptor to function in vivo as a homodimer or a heterodimer is unknown.

Thus, in accordance with another embodiment of the present invention, there is provided a method to render a steroid/thyroid receptor protein capable of forming a homodimer, said method comprising introducing the DNA binding domain of GAL4 into said receptor protein.

As employed herein, the term "ligand or ligand precursor for a member of the steroid/thyroid receptor protein superfamily" (i.e., intracellular receptor) refers to a substance or compound which (in its unmodified form, 15 or after conversion to its "active" form), inside a cell, binds to the receptor protein, thereby ligand/receptor complex, which in turn can activate an appropriate hormone response element. A ligand therefore is a compound which acts to modulate gene transcription for 20 a gene maintained under the control of a hormone response element, and includes compounds such as hormones, growth substances, non-hormone compounds that modulate growth, and the like. Ligands include steroid or steroid-like hormone, retinoids, thyroid hormones, pharmaceutically 25 compounds, and the like. Individual ligands may have the ability to bind to multiple receptors.

Accordingly, as employed herein, "putative ligand" refers to compounds such as steroid or steroid-like hormones, pharmaceutically active compounds, and the like, which are suspected to have the ability to bind to the orphan receptor of interest, and to modulate transcription of genes maintained under the control of response elements recognized by such orphan receptor.

Examples of known ligands include all-transretinoic acid (ligand for retinoic acid receptor), 9-cisretinoic acid (ligand for retinoid X receptor),
dexamethasone (ligand for glucocorticoid receptor), thyroid
hormone (ligand for thyroid hormone receptor), 1,25dihydroxy vitamin D₃ (ligand for vitamin D₃ receptor), and
the like.

Examples of known ligand precursors include alltrans-retinoic acid (which can be converted to 9-cisretinoic acid), 25-hydroxy vitamin D₃ (which can be
converted to 1,25-dihydroxy vitamin D₃), \$\beta\$-carotene,
biologically active lipids (e.g., fatty acids such as
linoleic acid or arachidonic acid), and the like.

As employed herein, the term "members of the 15 steroid/thyroid receptor protein superfamily" (also known as "nuclear receptors" or "intracellular receptors") refers to hormone binding proteins that operate as liganddependent transcription factors. Members of the steroid/thyroid receptor protein superfamily for which 20 specific ligands have not yet been identified are referred to herein as "orphan receptors". These hormone binding proteins have the intrinsic ability to bind to specific DNA sequences. Following binding, the ligand/orphan receptor complex functions to modulate the transcriptional activity 25 of target gene (i.e., a gene associated with the specific DNA sequence).

It is useful to distinguish the terms receptor "subtype" and receptor "class". For example, retinoid responsive receptors comprise a "class" of receptors, all of which are responsive to retinoid compounds. Similarly, thyroid hormone receptors comprise a "class" of receptors which are responsive to thyroid hormone. Each class can be divided into various subtypes, i.e., specific members of the class which have different tissue distributions,

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different affinities for the native ligand, different activation properties when contacted with the native ligand, and so on.

Some classes of receptors include sub-families of 5 distinctly different types of receptors. example, while the retinoid class of receptors includes both the retinoic acid receptors (RARs) and the retinoid X receptors (RXRs), these two different sub-families are clearly distinct. For example, each member of the RAR 10 sub-family is responsive to a defined first hormone response element (HRE), and each member of the RXR sub-family is responsive to a defined second HRE (which is distinctly different from the first HRE). Accordingly, in accordance with one aspect of the present invention, there is provided a method to identify compounds which are selective for a specific steroid/thyroid receptor protein from among a plurality of such receptors which respond to a common ligand (i.e., different subtypes of the same subfamily or members of the same class, but a different sub-20 family), said method comprising

introducing the DNA binding domain of GAL4 into each one of said plurality of receptor proteins, thereby producing a plurality of modified receptor proteins,

contacting cells containing one of said modified receptor proteins with a putative selective ligand for said modified receptor protein;

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wherein said cells contain a GAL4 response element operatively linked to a reporter gene,

monitoring the expression of reporter gene product, then

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repeating said contacting and monitoring steps with a different modified receptor protein until all members of said plurality of modified receptor proteins have individually undergone said contacting, and thereafter

identifying those compounds which are selective for one specific steroid/thyroid receptor protein from among said plurality of receptors which respond to a common ligand.

There are a number of receptors which respond to a common ligand, thus the plurality of receptors contemplated by the above-described method can be selected from the multiple subtypes of a receptor responsive to a common ligand, e.g., the various subtypes of RAR and/or the various subtypes of RXR, the various subtypes of PPAR, the various subtypes of COUP, and the like.

Such compounds are useful, for example, for the treatment of steroid or steroid-like hormone-responsive disease states. As used herein, the phrase "steroid or steroid-like hormone responsive disease state" refers to:

- (i) any disease state wherein a gene product (or a portion of a gene product) not normally subject to steroid or steroid-like hormone expression control is placed, by translocation, under the control of a steroid or steroid-like hormone responsive sequence, or
- (ii) any disease state wherein a first gene product (or a portion of a gene product) subject to steroid or steroid-like hormone expression control by a first steroid or steroidlike hormone is placed, by translocation, under the control of a second steroid or steroid-like hormone responsive sequence, or

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- (iii) any disease state which correlates with the expression of abnormal gene product, wherein said gene product (or a portion of said gene product) is normally subject to steroid or steroid-like hormone expression control, or
- (iv) any disease state which correlates with an abnormal level of expression of gene product, the expression of which is normally maintained under steroid or steroid-like hormone expression control, or
 - (v) any disease state which correlates with an abnormal level of receptor, the presence of which is normally maintained under steroid or steroidlike hormone expression control, or
- (vi) any disease state which correlates with an abnormal level of ligand, the presence of which is normally maintained under steroid or steroidlike hormone expression control.

As employed herein, the phrase "compounds which 20 are selective for a specific steroid/thyroid receptor protein from among a plurality of such receptors which respond to a common ligand" refers to compounds which interact with the receptor subtype associated with said steroid or steroid-like hormone responsive disease state to 25 a significantly greater extent than with other subtypes or sub-families of the same receptor class, i.e., compounds which are preferentially selective for one receptor subtype (or sub-family) in modulating the transcription activation properties thereof. The terminology "significantly greater 30 extent", as applied to interaction between ligand and a specific receptor subtype (or sub-family), refers to ligands which have a significantly higher therapeutic index (i.e., the ratio of efficacy to toxicity) for treatment of the target disease state than for activation of pathways 35 mediated by other subtypes (or sub-families) of the same receptor class. The toxicity of therapeutic compounds

frequently arises from the non-selective interaction of the therapeutic compound with receptor subtypes (or subfamilies) other than the desired receptor subtype (or subfamily). Thus, the present invention provides a means to dramatically reduce the incidence of side-reactions commonly associated with hormone therapy.

The DNA-binding domains of all members of the steroid/thyroid receptor protein superfamily are related, consisting of 66-68 amino acid residues, and possessing about 20 invariant amino acid residues, including nine cysteines.

A member of the superfamily can be identified as a protein which contains the above-mentioned invariant amino acid residues, which are part of the DNA-binding domain of such known steroid receptors as the human glucocorticoid receptor (amino acids 421-486), the estrogen receptor (amino acids 185-250), the mineralocorticoid receptor (amino acids 603-668), the human retinoic acid receptor α1 (amino acids 88-153). The highly conserved amino acids of the DNA-binding domain of members of the superfamily are as follows:

wherein X designates non-conserved amino acids within the DNA-binding domain; the amino acid residues denoted with an

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asterisk are residues that are almost universally conserved, but for which variations have been found in some identified hormone receptors; and the residues enclosed in parenthesis are optional residues (thus, the DNA-binding domain is a minimum of 66 amino acids in length, but can contain several additional residues).

Exemplary members of the steroid/thyroid superfamily of receptors include steroid receptors such as glucocorticoid receptor, mineralocorticoid receptor, 10 progesterone receptor, androgen receptor, vitamin D. receptor, and the like; plus retinoid receptors, such as RAR α , RAR β , RAR γ , and the like, plus RXR α , RXR β , RXR γ , and the like; thyroid receptors, such as $TR\alpha$, $TR\beta$, and the like; as well as other gene products which, by their 15 structure and properties, are considered to be members of the superfamily, as defined hereinabove. Examples of orphan receptors include HNF4 (see, for example, Sladek et al., in Genes & Development 4: 2353-2365 (1990)], the COUP family of receptors [see, for example, Miyajima et al., in 20 Nucleic Acids Research <u>16</u>: 11057-11074 (1988), Wang et al., in Nature <u>340</u>: 163-166 (1989), including the COUP-like receptors and COUP homologs, such as those described by Mlodzik et al., in Cell <u>60</u>: 211-224 (1990) and Ladias et al., in Science 251: 561-565 (1991)], the ultraspiracle 25 receptor [see, for example, Oro et al., in Nature 347: 298-301 (1990)], PPAR [see, for example, Dreyer et al., in Cell 68:879-887 (1992)], and the like.

As set forth herein, the DNA binding domain of GAL4 protein is utilized in the present invention in the identification of ligands or ligand-precursors for orphan receptors; and in the identification of compounds selective for a specific steroid/thyroid receptor protein from among a plurality of steroid/thyroid receptor proteins. The DNA binding domain of the yeast GAL4 protein comprises at least the first 74 amino acids thereof (see, for example, Keegan

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et al., Science 231:699-704 (1986)). Preferably, the first 90 or more amino acids of the GAL4 protein will be used, with the first 147 amino acid residues of yeast GAL4 being presently most preferred.

5 The GAL4 fragment employed in the practice of the present invention can be incorporated into any of a number of sites within a steroid/thyroid receptor protein. example, the GAL4 DNA binding domain can be introduced at the amino terminus of a steroid/thyroid receptor protein, 10 or the GAL4 DNA binding domain can be substituted for the native DNA binding domain of a steroid/thyroid receptor, or the GAL4 DNA binding domain can be introduced at the carboxy terminus of a steroid/thyroid receptor protein, or at other positions as can readily be determined by those of 15 skill in the art. Thus, for example, a modified receptor protein can be prepared which consists essentially of amino acid residues 1-147 of GAL4, plus the ligand binding domain of said steroid/thyroid receptor protein (i.e., containing the ligand binding domain only of said receptor, 20 substantially absent the DNA binding domain and amino terminal domain thereof).

Identification methods according to the present invention involve the use of a functional bioassay system, wherein the modified receptor and a reporter plasmid are cultured in suitable host cells in the presence of test compound. Evidence of transcription (e.g., expression) of reporter gene is then monitored to determine the presence of an activated receptor-ligand complex. Accordingly, the functional bioassay system utilizes two plasmids: an "expression" plasmid and a "reporter" plasmid. The expression plasmid can be any plasmid which contains and is capable of expressing DNA encoding the desired modified form of orphan steroid/thyroid receptor protein, in a suitable host cell. The reporter plasmid can be any

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plasmid which contains an operative GAL4 response element functionally linked to an operative reporter gene.

Exemplary GAL4 response elements are those containing the palindromic 17-mer:

5 5'-CGGAGGACTGTCCTCCG-3' (SEQ ID NO:5),

such as, for example, 17MX, as described by Webster et al., in Cell <u>52</u>:169-178 (1988), as well as derivatives thereof. Additional examples of suitable response elements include those described by Hollenberg and Evans in Cell <u>55</u>:899-906 (1988); or Webster et al. in Cell <u>54</u>:199-207 (1988).

Exemplary reporter genes include chloramphenicol transferase (CAT), luciferase (LUC), beta-galactosidase $(\beta$ -gal), and the like. Exemplary promoters include the simian virus (SV) promoter or modified form thereof (e.g., ΔSV), the thymidine kinase (TK) promoter, the mammary tumor virus (MTV) promoter or modified form thereof (e.g., AMTV), and the like [see, for example, Mangelsdorf et al., in Nature 345:224-229 (1990), Mangelsdorf et al., in Cell 66:555-561 (1991), and Berger et al., in J. Steroid 20 Biochem. Molec. Biol. <u>41</u>:733-738 (1992)]. The plasmids pGMCAT, pGHCAT, ΔMTV-GAL_D3-LUC, pTK-GAL_o3-LUC, AMTV-GAL, 3-CAT, and the like, are examples of reporter plasmids which contain an operative hormone responsive promoter/enhancer element functionally linked 25 operative reporter gene, and can therefore be used in the above-described functional bioassay (see Example 2 for details on the preparation of these plasmids). the operative hormone responsive promoter/enhancer element is the MTV LTR; in pGHCAT it is the functional portion of 30 the growth hormone promoter. In both pGMCAT and GHCAT the operative reporter gene is the bacterial gene chloramphenicol acetyltransferase (CAT).

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As used herein in the phrase "operative response element functionally linked to an operative reporter gene", the word "operative" means that the respective DNA sequences (represented by the terms "GAL4 response element" 5 and "reporter gene") are operational, i.e., work for their intended purposes; the word "functionally" means that after the two segments are linked, upon appropriate activation by a ligand-receptor complex, the reporter gene will be expressed as the result of the fact that the "GAL4 response element" was "turned on" or otherwise activated.

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In practicing the above-described functional bioassay, the expression plasmid and the reporter plasmid are co-transfected into suitable host cells. transfected host cells are then cultured in the presence 15 and absence of a test compound to determine if the test compound is able to produce activation of the promoter operatively linked to the GAL4 response element of the reporter plasmid. Thereafter, the transfected and cultured host cells are monitored for induction (i.e., the presence) 20 of the product of the reporter gene sequence.

Any cell line can be used as a suitable "host" for the functional bioassay contemplated for use in the practice of the present invention. Thus, in contrast to the requirements of prior art assay systems, there is no need to use receptor-negative cells in carrying out the invention process. Since the modified receptor employed in the practice of the present invention is the only species test cell which is capable of initiating transcription from a GAL4 response element, the expression 30 of native receptor by the test cell does not contribute to background levels. Thus, the invention bioassay is very selective.

Cells contemplated for use in the practice of the present invention include transformed cells,

transformed cells, neoplastic cells, primary cultures of different cell types, and the like. Exemplary cells which can be employed in the practice of the present invention include Schneider cells, CV-1 cells, HuTu80 cells, F9 5 cells, NTERA2 cells, NB4 cells, HL-60 cells, 293 cells, Hela cells, yeast cells, and the like. Preferred host cells for use in the functional bioassay system are COS cells and CV-1 cells. COS-1 (referred to as COS) cells are monkey kidney cells that express SV40 T antigen (Tag); 10 while CV-1 cells do not express SV40 Tag. The presence of Tag in the COS-1 derivative lines allows the introduced expression plasmid to replicate and provides a relative increase in the amount of receptor produced during the assay period. CV-1 cells are presently preferred because 15 they are particularly convenient for gene transfer studies and provide a sensitive and well-described host cell system.

The above-described cells (or fractions thereof) are maintained under physiological conditions when contacted with physiologically active compound. "Physiological conditions" are readily understood by those of skill in the art to comprise an isotonic, aqueous nutrient medium at a temperature of about 37°C.

In accordance with another embodiment of the present invention, there is provided a method of screening for antagonists of steroid/thyroid receptor proteins, said method comprising

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culturing test cells containing

(i) increasing concentrations of at least one compound whose ability to inhibit the transcription activation activity of hormone receptor agonists is sought to be determined, and

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(ii) optionally, at least one agonist
for said steroid/thyroid receptor protein(s)
or functional modified forms thereof,

wherein said test cells contain

(i) exogenous DNA which expresses a modified form of a steroid/thyroid receptor protein(s) or functional modified forms thereof, wherein the modified form of the steroid/thyroid receptor protein(s) or functional modified forms thereof contains the DNA binding domain of GAL4, and

(ii) a GAL4 response element operatively linked to a reporter gene; and thereafter

assaying for evidence of transcription of said reporter gene in said cells as a function of the concentration of said compound in said culture medium, thereby indicating the ability of said compound to inhibit activation of transcription by hormone receptor agonists.

Media employed for such culturing may include agonist for the receptor being tested, or the receptor may be constitutive (i.e., not require the presence of agonist for activation), or a fixed concentration of agonist can be added to the media employed for such testing.

The above-described assays of the present invention have low background and a broad dynamic range.

In accordance with yet another embodiment of the 30 present invention, there is provided a method for identifying biologically active compounds characterized by being:

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convertible into ligand or ligand precursor for intracellular receptor(s), or

capable of effecting transformation of a second compound into ligand or ligand precursor for intracellular receptor(s), or

capable of effecting transformation of an intracellular receptor into its active form,

said method comprising:

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- (1) contacting cells or cell fractions with said biologically active compound under physiological conditions;
 - (2) fractionating the cells, cell fractions and/or the medium in which said contacting is carried out; and
- 15 (3) subjecting each of said fractions to a functional assay wherein each fraction is cultured with host cells containing DNA encoding the above-described GAL4-receptor chimeric constructs, and a GAL4 response element operatively linked to reporterencoded DNA, and expression of reporter gene is monitored to determine the presence of an activated receptor-ligand complex.

Examples of ligands and ligand precursors for intracellular receptor(s) have been set forth above. Compounds capable of effecting transformation of an intracellular receptor into its active form include compounds which act as activators of receptors by alternative biochemical pathways not involving direct interaction of the compound with the receptor. Examples of such compounds include dopamine, fatty acids, clofibric acid, xenobiotics, and the like.

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A wide variety of cells are contemplated for use in the practice of this embodiment of the present invention, as noted above with respect to alternate embodiments of the invention.

5 Cell fractions contemplated for use in this embodiment of the present invention include a variety of subcellular fractions, e.g., nuclear fractions, cytosolic fractions, membrane fractions, and the like.

Fractionating the cells, cell fractions and/or medium in which the above-described contacting is carried out can be conducted in a variety of ways, as can be readily determined by those of skill in the art. For example, the fermentation broth can be extracted into aqueous and/or organic media and further fractionated and purified by a variety of physical separation techniques, such as, for example, standard column chromatography, high performance liquid chromatography (HPLC) separation, ultracentrifugation (e.g., density gradient or isopycnic banding), thin layer chromatography (TLC), and the like.

20 The invention identification method involves the use of a functional bioassay system, wherein each of the above-described fraction(s) is cultured with host cells containing DNA encoding the above-described GAL4-receptor chimeric constructs, and GAL4 response element 25 operatively linked to reporter-encoded DNA, and expression of reporter gene product is monitored to determine the presence of an activated receptor-ligand Accordingly, the functional bioassay system utilizes two plasmids: an "expression" plasmid and a "reporter" plasmid. The expression plasmid can be any plasmid which contains and is capable of expressing the above-described GAL4receptor chimeric constructs (or mutants thereof), in a suitable host cell. The reporter plasmid can be any plasmid which contains an operative GAL4

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promoter/enhancer element functionally linked to an operative reporter gene.

Exemplary reporter genes suitable for use in the practice of this embodiment of the present invention are as described above with respect to alternate embodiments of the invention.

When using the functional bioassay system to determine the functionality of ligands for intracellular receptors, i.e., members of the steroid/thyroid superfamily of receptors, in preferred forms, plasmids will carry a selectable marker such as the amp gene. In addition, in preferred forms the reporter plasmids will have a suitable hormone responsive promoter/enhancer element e.g., the MTV LTR or a functional portion of the growth hormone promoter.

- Expression plasmids containing the SV40 origin of replication (ori) can propagate to high copy number in any host cell which expresses SV40 Tag. Thus expression plasmids carrying the SV40 "ori" can replicate in COS cells, but not in CV-1 cells. Although the increased expression afforded by high copy number is desired, it is not critical to the operability of the "cis-trans" bioassay system. The expression vectors typically employed in the bioassay are generally so efficient that the assay works in virtually any host.
- Thus, as described above with respect to other embodiments of the present invention, any cell line can be used as a suitable "host" for the functional bioassay contemplated for use in this embodiment of the present invention.
- Based on the above-described functional bioassays, the activation of receptor by steroids or steroid-like compounds (or the inhibition of such

activation by test compounds in the presence of agonists) can be examined.

In accordance with still another embodiment of the present invention, there is provided a method for identifying cell lines which are capable of promoting the conversion of biologically active compounds into:

- (A) ligand or ligand precursor for intracellular receptor(s), or
- (B) species which promote the transformation of a second compound into ligand or ligand precursor for intracellular receptor(s), or
- (C) species which are capable of effecting transformation of an intracellular receptor into its active form,
- 15 said method comprising:

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- (1) contacting cells with biologically active compound(s) under conditions suitable to maintain the viability of said cells;
- (2) fractionating the cells and the medium in which the cells are maintained;
 - (3) subjecting each of said fractions to a functional bioassay wherein each fraction is cultured with host cells containing DNA encoding the above-described GAL4-receptor chimeric constructs, and a GAL4 response element operatively linked to reporter-encoded DNA, and expression of reporter gene is monitored to determine the presence of an activated receptor-ligand complex; and
- (4) identifying those cell lines which promote the conversion of said biologically active compounds into products functional in said functional bioassay.

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In accordance with a still further embodiment of the present invention, there is provided a method for identifying enzymatic or non-enzymatic activities capable of promoting the conversion of biologically active 5 compounds into:

- (A) ligand or ligand precursor for intracellular receptor(s), or
- (B) species which promote the transformation of a second compound into ligand or ligand precursor for intracellular receptor(s), or
- (C) species which are capable of effecting transformation of an intracellular receptor into its active form,

said method comprising:

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- 15 (1) subjecting biologically active compound(s) to conditions suitable to promote the desired conversion;
 - (2) subjecting the reaction product(s) of step (1) to a functional bioassay wherein each fraction is cultured with host cells containing DNA encoding the above-described GAL4-receptor chimeric constructs, and a GAL4 response element operatively linked to reporter-encoded DNA, and expression of reporter gene is monitored to determine the presence of an activated receptor-ligand complex; and
 - (3) identifying those conditions which promote the conversion of said biologically active compounds into products functional in said functional assay as providing the desired activities.

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Conditions suitable to promote the desired conversion of biologically active compound include contacting with cell fractions which potentially contain the desired activity, subjecting the biologically active compound to temperature variations, pH changes, ultraviolet radiation, or any other chemical or physical exposure which has the potential to effect a modification of the activity of the biologically active compound.

The invention will now be described in greater detail by reference to the following non-limiting examples.

Example 1 Preparation of GAL4-receptor fusion proteins

A basic vector useful for the generation of GAL4receptor fusion proteins is called pCMX-GAL4 (see SEQ ID
15 NO:2). This vector encodes GAL4 DNA binding domain,
followed by a polylinker sequence useful in the cloning.
The parental expression vector pCMX has been described by
Umesono et al., in Cell 65:1255-1266 (1991), and the GAL4
portion of pCMX-GAL4 is derived from plasmid pSG424,
20 described by Sadowski and Ptashne, in Nucleic Acids Res.
17:7539 (1989).

In general, GAL4-receptor ligand binding domain fusions are prepared by taking advantage of mutant receptor cDNA clones, such as GR-RAR chimera [see, for example, Giguere et al., in Nature 330:624-629 (1987)]. These mutant receptor cDNAs encode common XhoI sites at the end of the DNA binding domain, as described by Giguere et al., supra. To do so, a new pCMX-GAL4 vector was prepared which encodes a compatible SalI site in the polylinker sequence (there is an XhoI site in the GAL4 sequence):

SalI site: G'TCGAC XhoI site: C'TCGAG

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This allows efficient transfer of the receptor ligand binding domain to GAL4 DNA binding domain. Through this method, a number of chimeric species have been generated, e.g., GAL4-hRARα, GAL4-hTRβ, GAL4-hGR, GAL4-hERR1, GAL4-hERR2, GAL4-hXR2 (hXR2 is described in co-pending application Serial No. 07/761,068, now pending), GAL4-mXR5 (mXR5 is described in co-pending application Serial No. 07/761,068, now pending), GAL4-hVDR, GAL4-hRXRα, and the like. For example, hRARα cDNA was mutated to encode a unique XhoI site as follows (see Giguere et al., supra):

AAA-GAG (K155-E156) - CTC-GAG (L155-E156)

This mutated RARa cDNA was then digested with XhoI and BamHI, and ligated into a SalI-BamHI digest of pCMX-GAL4 (containing the "new" linker set forth in SEQ ID NO:4, wherein the linker set forth in SEQ ID NO:4 is inserted into the construct set forth in SEQ ID NO:2 in place of residues 479-532 thereof), to produce a construct having the sequence, in relevant part:

GAL4 DBD---CCG-GAA-TTC-GGT-ACC-GTC-GAG-TCT-GTG---RAR LBD,

20 wherein "DBD" refers to DNA binding domain, and "LBD" refers to ligand binding domain.

If mutants of the type referred to above are not available the construction of GAL4-containing chimerics, one may simply look for any convenient 25 restriction enzyme site downstream of the DNA binding domain of the receptor of interest (i.e., within about the first 30 amino acid residues downstream of the conserved Gly-Met residues of the DNA binding domain, i.e., within 30 residues of the last two residues shown in SEQ ID NO:1), 30 and utilize the carboxy terminal sequences therefrom.

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Alternatively, one can utilize PCR-based mutagenesis to add any desired restriction sites to a segment of DNA. For example, primers can be designed which are complementary to the ligand binding domain of any 5 member of the steroid/thyroid superfamily of receptors. Such primers are capable of amplifying desired portions of a given receptor utilizing PCR. By adding the desired restriction sites to the ends of the primers, one can ensure that each copy synthesized by PCR will possess such 10 restriction enzyme sites at its termini [Scharf, Cloning with PCR. In: PCR Protocols (Innis et al., eds.) Academic Press Inc. (San Diego, CA) 84-91 (1990)]. Subsequently, the fragment can be digested with both enzymes and ligated into pCMX-GAL4, which has been digested with the same or 15 compatible enzymes.

Thus, for example, the ligand binding domain of chicken RARB can be amplified using PCR with primers complementary to amino acids 147-154 and 447-452. A SalI site can be added to the 5' end and a NheI site to the 3' end. After amplification, the fragment can be digested with SalI and NheI, then ligated to pCMX-GAL4 (which has been digested with SalI and NheI; employing pCMX-GAL4n which contains the polylinker set forth in SEQ ID NO:4).

Example 2

25 <u>Preparation of reporter constructs</u>

Various reporter constructs are used in the examples which follow. They are prepared as follows:

TK-LUC: The MTV-LTR promoter sequence was removed from the MTV-LUC plasmid described by Hollenberg and Evans in Cell <u>55</u>:899-906 (1988) by *HindIII* and *XhoI* digest, and cloned with the *HindIII-XhoI* fragment of the Herpes simplex virus thymidine kinase gene promoter (-105 to +51 with respect to the transcription start site, m,

isolated from plasmid pBLCAT2, described by Luckow & Schutz in Nucleic Acids Res. <u>15</u>:5490 (1987)) to generate parental construct TK-LUC.

pTK-TREp2-LUC: Two copies of double-stranded 5 palindromic thyroid hormone response element (TREp) oligonucleotides, encoding a retinoic acid receptor (RAR) binding site, were cloned upstream of the TK promoter of TK-LUC at the HindIII site.

pTK-GALp3-LUC: Three copies of double-stranded 10 17MX oligonucleotides, encoding a GAL4 binding site, were cloned upstream of the TK promoter of TK-LUC at the HindIII site.

AMTV-GALp3-CAT: Three copies of double-stranded 17MX oligonucleotides, encoding a GAL4 binding site, were cloned at the *Hin*dIII site in the AMTV-CAT plasmid described by Hollenberg and Evans, supra.

AMTV-GALp3-LUC: Three copies of double-stranded 17MX oligonucleotides, encoding a GAL4 binding site, were cloned at the *HindIII* site in the AMTV-LUC plasmid described by Hollenberg and Evans, <u>supra</u>. The AMTV-LUC plasmid encodes a mutant MTV-LTR promoter which no longer responds to hormones, as described by Hollenberg and Evans, <u>supra</u>.

CMX-BGAL: The coding sequence for the *E. coli*8-galactosidase gene was isolated from plasmid pCH110 [see
Hall et al., J. Mol. Appl. Genet. 2:101-109 (1983)] by
HindIII and BamHI digest, and cloned into pCMX eucaryotic
expression vector [see Umesono et al., supra].

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Example 3

Screening assay for receptor selective agonists

CV-1 cells are co-transfected with CMX-GAL-hRARα and tk-gal_px3-luc in 96 well plates. The usual amounts of DNA per 96 wells are 1 μg of CMX-GAL-hRARα, 5 μg of tk-gal_px3-luc, and 4 μg of CMX-BGAL. Typically, transfections are performed in triplicate. The plates are then incubated overnight at 37°C.

The cells are washed twice with fresh medium.

10 Fresh medium containing one concentration of a serial dilution of agonist is added to each well. A typical agonist dilution series extends from 10⁻⁵M through 10⁻¹¹M. A solvent control is performed for each agonist. The cells are incubated at 37°C for 1-2 days.

The cells are rinsed twice with buffered saline solution. Subsequently, cells are lysed, in situ, by adding 200 μ l of lysis buffer. After 30 minutes incubation at room temperature, 40 μ l aliquots of cell lysate are transferred to 96-well plates for luciferase reporter gene assays and 8-galactosidase transfection controls [see Heyman et al., Cell <u>68</u>:397-406 (1992)].

The data are expressed as relative light units (RLUs) per O.D. unit of \$\beta\$-galactosidase per minute. The triplicates are averaged for each concentration and plotted as normalized RLUs against the dose of agonist or as fold induction vs the dose of agonist.

The selectivity of an agonist for a particular receptor is measured by comparing the activation of that receptor with the activation seen by the same agonist utilizing other related receptors. It is frequently useful to express the activation elicited by a particular agonist as a percent of the activation elicited by the maximum

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concentration of all-trans retinoic acid on the same receptor.

This example demonstrates the ease with which the process of the present invention can be adapted for screening large numbers of compounds and/or cell lines to determine the presence of ligand or the ability of a test cell line to produce a functional ligand from the components of the media.

Example 4

Dose response of various hRARa constructs

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Effector plasmid, reporter plasmid, B-galactosidase control plasmid are co-transfected into CV-1 cells at a ratio of about 1:5.4, using a liposomemediated method, employing DOTAP (Boehringer Manheim) 15 according to manufacturer's instructions in serum-free media (e.g., Opti-MEM, from Life Technologies, Gaithersberg, MD). After about 18 hours, the cells are washed twice with fresh Opti-MEM and all-trans retinoic acid is added to the media to the final molar concentration 20 indicated in Figure 3. After 24 hours of incubation, the cells are rinsed with phosphate buffered saline (pH 7.2) and lysed. Aliquots are assayed for luciferase and B-galactosidase activity. Luciferase activity normalized to optical density units of B-galactosidase per 25 minute of incubation.

The data are expressed in Figure 3 as fold induction over the same construct incubated in solvent alone. The receptor:reporter combinations employed are as follows:

CMX-GAL-hRARα1 (wherein the ligand binding domain of hRARα is fused to amino acids 1-173 of GAL4) with pTK-GALp3-LUC;

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RS-GGR (wherein the amino-terminal and DNA binding domains of human glucocorticoid receptor alpha are fused to the ligand binding domain of hRARa) with MTV-LUC;

RS-GRR (wherein the amino-terminal domain of human glucocorticoid receptor alpha is fused to the DNA binding and ligand binding domains of hRARa) with pTK-TREp2-LUC; and CMX-hRARal with pTK-TREp2-LUC.

Review of Figure 3 reveals that chimeric receptors of the invention are much more responsive in agonist-type assays than are unmodified receptors, or chimeric receptors employed in the prior art (e.g., GRR or GGR).

While the invention has been described in detail with reference to certain preferred embodiments thereof, it will be understood that modifications and variations are within the spirit and scope of that which is described and claimed.

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Summary of Sequences

Sequence ID No. 1 is the consensus amino acid sequence of the DNA binding domain of members of the steroid/thyroid superfamily of receptors.

- Sequence ID No. 2 is a nucleic acid sequence (and the deduced amino acid sequence) of a vector useful for the preparation of GAL4-receptor fusion proteins, including an exemplary segment encoding a GAL4 DNA binding domain useful in the practice of the present invention.
- Sequence ID No. 3 is the deduced amino acid sequence of the nucleic acid sequence of SEQ ID NO:2.

Sequence ID No. 4 is the nucleic acid sequence of a linker useful in the preparation of chimeric constructs of the invention.

Sequence ID No. 5 is the nucleic acid sequence of a GAL4 response element, useful in the practice of the present invention.

PCT/US94/14426 WO 95/18380

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SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: Evans, Ronald M. Umesono, Kazuhiko Blumberg, Bruce Rangarajan, Pundi N.
- (ii) TITLE OF INVENTION: NOVEL USES FOR GAL4-RECEPTOR CONSTRUCTS
 - (iii) NUMBER OF SEQUENCES: 5
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Pretty, Schroeder, Brueggemann & Clark
 (B) STREET: 444 South Flower Street, Suite 2000

 - (C) CITY: Los Angeles
 - (D) STATE: California
 - (E) COUNTRY: USA (F) ZIP: 90071

 - (v) COMPUTER READABLE FORM:
 (A) MEDIUM TYPE: Floppy disk

 - (B) COMPUTER: IBM PC compatible
 (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 08/177,740
 - (B) FILING DATE: 30-DEC-1993
 (C) CLASSIFICATION:
 - (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Reiter, Stephen E.
 - (B) REGISTRATION NUMBER: 31,192
 - (C) REFERENCE/DOCKET NUMBER: P41 9392
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: 619-546-4737 (B) TELEFAX: 619-546-9392
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 71 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (v) FRAGMENT TYPE: N-terminal
 - (ix) FEATURE:
 - (A) NAME/KEY: Modified-site
 - (B) LOCATION: group (37..38, 45..47)
 - (D) OTHER INFORMATION: /label= Option /note= "These residues are optional. Thus, the DNA-binding domain is a minimum of 66 amino acids in length, but can contain several additional residues."

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	(ix	(FEATURE: (A) NAME/KEY: Region (B) LOCATION: one-of(7, 9, 11, 13, 22, 27, 62, 65, 70) (D) OTHER INFORMATION: /label= Conserved /note= "These residues are almost universally conserved, but variations have been found in some identified hormone receptors."														
	(xi) SE	QUEN	CE DE	escr:	[PTI	ON: S	EQ	ID N	0:1:				•			
	Су: 1	s Xa	a Xa	a Cys	Xaa 5	а Хаа	a Asr	ха	a Al	а Ха 10		y Xaa	Tyr	Xaá	Xaa 15	Xaa	
	Xa	а Су	s Xa	a Xaa 20	су:	s Ly	s Xaa	Ph	e Pb 25		a Ar	y Xaa	Xaa	Xaa 30	Xaa	Xaa	
	Xa	а Ха	a Xa: 35	a Xaa	Xaa	а Хаа	a Cys	Xa:	a Xa	a Xa	a Xaa	a Xaa	Xaa 45	Xaa	Xaa	Cys	
	Xa	a Xa 50		a Lys	: Xaa	a Xaa	a Arg 55	Xaa	a Xa	a Cy	s Xaa	Xaa 60	Cys	Arg	Xaa	Xaa	
	Ly: 65	s Су	s Xaa	a Xaa	Xaa	Gl ₃ 70	/ Met										
(2)	INF	ORMA'	TION	FOR	SEQ	ID 1	₹O:2:										
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 546 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: both																
	(ii)	MOI	LECUI	E TY	PE:	DNA	(gen	omi.	=)								
	(iii)	ну	POTHI	TICA	L: N	ro	•										
	(iv)	AN	ri-se	ense :	NO												
	(ix)	(2		e: Me/k Cati			ı (35 .	. 532	2, 5:	36!	544)						
	(xi)	SEC	OUENC	E DE	SCRI	PTIC	n: s	EQ 1	D N	0:2:							
GGG2	AGACC	CA I	\GCT1	GAAG	C AA	.GCCI	CCTG	AAA				A CTO		Ser		52	2
												CTC # Leu I				100	
_	_					_						AAC T Asn T 35		_		148	3
_					_							AGG G Arg A	_			196	5

ACA GAA GTG GAA TCA AGG CTA GAA AGA CTG GAA CAG CTA TTT CTA CTG
Thr Glu Val Glu Ser Arg Leu Glu Arg Leu Glu Gln Leu Phe Leu Leu
55 60 65

33

					GAC Asp			 	 TTA Leu	292
					ACA Thr					340
					AGA Arg 110					388
					AGA Arg				 	436
					AGA Arg					484
					CAG Gln		 	 	 	532
TAG	GTA Val	 	GG							546

(2) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 169 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

Met Lys Leu Leu Ser Ser Ile Glu Gln Ala Cys Asp Ile Cys Arg Leu
1 10 15

Lys Lys Leu Lys Cys Ser Lys Glu Lys Pro Lys Cys Ala Lys Cys Leu 20 25 30

Lys Asn Asn Trp Glu Cys Arg Tyr Ser Pro Lys Thr Lys Arg Ser Pro

Leu Thr Arg Ala His Leu Thr Glu Val Glu Ser Arg Leu Glu Arg Leu 50 55

Glu Gln Leu Phe Leu Leu Ile Phe Pro Arg Glu Asp Leu Asp Met Ile 65 70 75 80

Leu Lys Met Asp Ser Leu Gln Asp Ile Lys Ala Leu Leu Thr Gly Leu 85 90 95

Phe Val Gln Asp Asn Val Asn Lys Asp Ala Val Thr Asp Arg Leu Ala 100 105 110

Ser Val Glu Thr Asp Met Pro Leu Thr Leu Arg Gln His Arg Ile Ser

Ala Thr Ser Ser Ser Glu Glu Ser Ser Asn Lys Gly Gln Arg Gln Leu 130 135 140

34

Thr Val Ser Pro Glu Phe Pro Gly Ile Arg Arg Arg Tyr Gln Ile Ser 145 150 155 160

Gly Ser Trp Pro Ala Ser Val Ala Arg 165

- (2) INFORMATION FOR SEQ ID NO:4:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 57 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (iii) HYPOTHETICAL: NO
 - (iv) ANTI-SENSE: NO
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

 CCGGAATTCG GTACCGTCGA CGATATCCCC CGGGCTGCAG GATCCTGGCC AGCTAGC 57
- (2) INFORMATION FOR SEQ ID NO:5:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 17 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: both
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (iii) HYPOTHETICAL: NO
 - (iv) ANTI-SENSE: NO
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:
 CGGAGGACTG TCCTCCG

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That which is claimed is:

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1. A method to determine the ligand or ligand precursor for an orphan member of the steroid/thyroid receptor protein superfamily, said method comprising

contacting cells containing a modified form of said orphan member with a putative ligand for said orphan member;

wherein said modified form of said orphan member is produced by introducing the DNA binding domain of GAL4 into said orphan member, and

wherein said cells contain a GAL4 response element operatively linked to a reporter gene, and thereafter

monitoring the expression of reporter gene product.

- 2. A method according to claim 1 wherein the DNA binding domain of GAL4 comprises amino acid residues 1-147 of GAL4.
- 3. A method according to claim 1 wherein the DNA binding domain of GAL4 is introduced at the amino terminus of the orphan member.
- 4. A method according to claim 1 wherein the DNA binding domain of GAL4 replaces the DNA binding domain of the orphan member.

- 5. A method to identify compounds which are selective for a specific steroid/thyroid receptor protein from among a plurality of such receptors which respond to a common ligand, said method comprising
- introducing the DNA binding domain of GAL4 into each one of said plurality of receptor proteins, thereby producing a plurality of modified receptor proteins,
- contacting cells containing one of said

 modified receptor proteins with a putative selective ligand for said modified receptor protein;

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wherein said cells contain a GAL4 response element operatively linked to a reporter gene,

monitoring the expression of reporter gene product, then

repeating said contacting and monitoring steps with a different modified receptor protein until all members of said plurality of modified receptor proteins have individually undergone said contacting, and thereafter

identifying those compounds which are selective for one specific steroid/thyroid receptor protein from among said plurality of receptors which respond to a common ligand.

6. A method according to claim 5 wherein said plurality of receptor proteins comprises multiple subtypes of a receptor responsive to a common ligand.

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- 7. A method according to claim 5 wherein the DNA binding domain of GAL4 comprises at least the first 74 amino terminal amino acid residues of GAL4.
- 8. A method according to claim 5 wherein the DNA binding domain of GAL4 is introduced at the amino terminus of the receptor protein.
- 9. A method according to claim 5 wherein the DNA binding domain of GAL4 replaces the DNA binding domain of the receptor protein.
- 10. A method according to claim 5 wherein said modified receptor protein consists essentially of amino acid residues 1-147 of GAL4, plus the ligand binding domain of said steroid/thyroid receptor protein.
- 11. A method of screening for antagonists of steroid/thyroid receptor proteins, said method comprising

culturing test cells containing

- (i) increasing concentrations of at least one compound whose ability to inhibit the transcription activation activity of hormone receptor agonists is sought to be determined, and
- (ii) optionally, at least one agonist
 for said steroid/thyroid receptor protein(s)
 or functional modified forms thereof,

wherein said test cells contain

(i) exogenous DNA which expresses a modified form of a steroid/thyroid receptor protein(s) or functional modified forms thereof, wherein the modified form of the steroid/thyroid receptor protein(s) or functional

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modified forms thereof contains the DNA binding domain of GAL4, and

(ii) a GAL4 response element operatively linked to a reporter gene; and thereafter

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assaying for evidence of transcription of said reporter gene in said cells as a function of the concentration of said compound in said culture medium, thereby indicating the ability of said compound to inhibit activation of transcription by hormone receptor agonists.

- 12. A method to render a steroid/thyroid receptor protein capable of forming a homodimer, said method comprising introducing the DNA binding domain of GAL4 into said receptor protein.
- 13. A method according to claim 12 wherein the DNA binding domain of GAL4 comprises at least the first 74 amino terminal amino acid residues of GAL4.
- 14. A method according to claim 12 wherein the DNA binding domain of GAL4 is introduced at the amino terminus of the receptor protein.
- 15. A method according to claim 12 wherein the DNA binding domain of GAL4 replaces the DNA binding domain of the receptor protein.
- 16. A method according to claim 12 wherein said modified receptor protein consists essentially of amino acid residues 1-147 of GAL4, plus the ligand binding domain of said steroid/thyroid receptor protein.

17. A method for identifying biologically active compounds characterized by being:

convertible into ligand or ligand precursor for intracellular receptor(s), or

capable of effecting transformation of a second compound into ligand or ligand precursor for intracellular receptor(s), or

capable of effecting transformation of an intracellular receptor into its active form,

10 said method comprising:

- contacting cells or cell fractions with said biologically active compound under physiological conditions;
- (2) fractionating the cells, cell fractions and/or the medium in which said contacting is carried out; and
 - (3) subjecting each of said fractions to a functional assay wherein each fraction is cultured with host cells containing DNA encoding a chimeric receptor comprising a member of the steroid/thyroid receptor protein superfamily having incorporated therein the DNA binding domain of the yeast GAL4 protein, and a GAL4 response element operatively linked to reporter-encoded DNA, and expression of reporter gene is monitored to determine the presence of an activated receptor-ligand complex.

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- 18. A method for identifying cell lines which are capable of promoting the conversion of biologically active compounds into:
 - (A) ligand or ligand precursor for intracellular receptor(s), or
 - (B) species which promote the transformation of a second compound into ligand or ligand precursor for intracellular receptor(s), or
 - (C) species which are capable of effecting transformation of an intracellular receptor into its active form,

said method comprising:

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- (1) contacting cells with biologically active compound(s) under conditions suitable to maintain the viability of said cells;
- (2) fractionating the cells and the medium in which the cells are maintained;
- (3) subjecting each of said fractions to a functional bioassay wherein each fraction is cultured with host cells containing DNA encoding a chimeric receptor comprising a member of the steroid/thyroid receptor protein superfamily having incorporated therein the DNA binding domain of the yeast GAL4 protein, and a GAL4 response element operatively linked to reporter-encoded DNA, and expression of reporter gene is monitored to determine the presence of an activated receptor-ligand complex; and
- (4) identifying those cell lines which promote the conversion of said biologically active compounds into products functional in said functional bioassay.

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- 19. A method for identifying enzymatic or nonenzymatic activities capable of promoting the conversion of biologically active compounds into:
 - (A) ligand or ligand precursor for intracellular receptor(s), or
 - (B) species which promote the transformation of a second compound into ligand or ligand precursor for intracellular receptor(s), or
 - (C) species which are capable of effecting transformation of an intracellular receptor into its active form,

said method comprising:

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- (1) subjecting biologically active compound(s) to conditions suitable to promote the desired conversion;
- subjecting the reaction product(s) of step (1) to a functional bioassay wherein each fraction is cultured with host cells containing DNA encoding a chimeric receptor comprising a member of the steroid/thyroid receptor protein superfamily having incorporated therein the DNA binding domain of the yeast GAL4 protein, and a GAL4 response element operatively linked reporter-encoded DNA, and expression of reporter gene is monitored to determine the presence of an activated receptor-ligand complex; and
- (3) identifying those conditions which promote the conversion of said biologically active compounds into products functional in said functional assay as providing the desired activities.

AAAAGCATTGTTAACAGGATTATTTGTACAAGATAATGTGAATAAAGATGCCGTCACAGA 109 4 Ω × Z > Z Ω O BsrGI U Hpal Ø 301

TAGATTGGCTTCAGTGGAGACTGATATGCCTCTAACATTGAGACAGCATAGAATAAGTGC 129 Ø S 出 Ø 吖 Ц H 口 Д Σ Д Н **(1)** S Ø Ц 361

2/4 GACATCATCGGAAGAGAGTAGTAACAAAGGTCAAAGACAGTTGACTGTATCGCCGGA 团 Д sedneuce ends--> Ø > H Ы Ø 2 GAL4 Ø Ö × Z ഗ ഗ 回 回 ഗ Ø ß

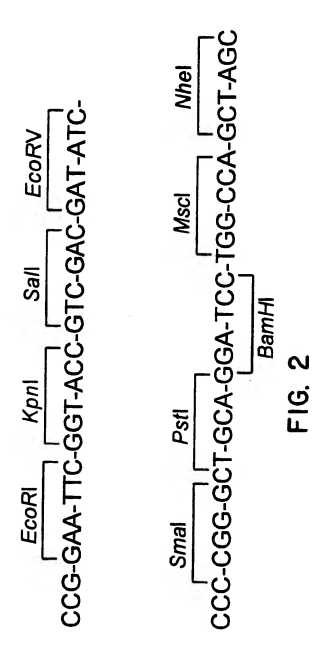
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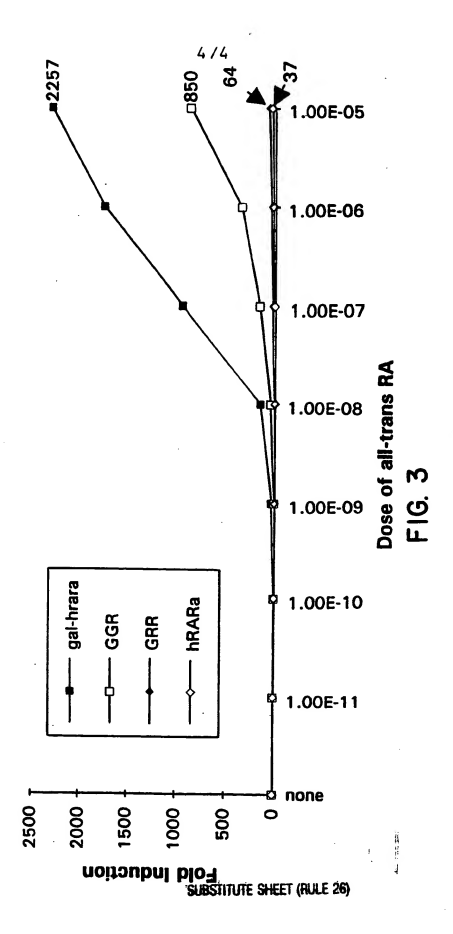
SUBSTITUTE SHEET (RULE 26)

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INTERNATIONAL SEARCH REPORT

ernational application No. PCT/US94/14426

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A. CLASSIFICATION OF SUBJECT MATTER IPC(6) :G01N 33/566; C12P 21/00; C12Q 1/00; C12N 15/00, 1/20 US CL :436/501; 435/69.1, 252.3, 7.2, 69.7, 172.3 According to International Patent Classification (IPC) or to both national classification and IPC						
B. FIELDS SEARCHED						
Minimum documentation searched (classification system followed by classification symbols)						
U.S. : 436/501; 435/69.1, 252.3, 7.2, 69.7, 172.3						
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched						
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) APS, DIALOG - Biotech Files						
C. DOCUMENTS CONSIDERED TO BE RELEVANT						
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.			
X	Proceedings of the National Academy of Sciences USA, 1-10 Volume 90, issued January 1993, G. Allenby et al, "Retinoic"					
Y	acid receptors and retinoid X reendogenous retinoic acids", pedocument.	11-19				
Y	Cell, Volume 54, issued 15 July al, "The Hormone Binding Doma Glucocorticoid Receptors Contain Activation Function", pages 199-2	ains of the Estrogen and an Inducible Transcription	1-19			
X Further documents are listed in the continuation of Box C. See patent family annex.						
Special categories of cited documents: To be document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention						
to be of particular relevance E* cartier document published on or after the international filing date "X" document of particular relevance; the claimed invention cannot be considered as ovel or cannot be considered as overland as overla						
cited	ument which may throw doubts on priority claim(s) or which is d to establish the publication date of another citation or other	when the document is taken alone 'Y' document of perticular colorance: the				
special reason (as specified) O document referring to an oral disclosure, use, exhibition or other means o document referring to an oral disclosure, use, exhibition or other means o document referring to an oral disclosure, use, exhibition or other means o document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art						
P* document published prior to the internstional filing date but later than "&" document member of the same patent family the priority date claimed						
Date of the actual completion of the international search Date of mailing of the international search report						
23 MARCH 1995 0 6 APR 1995						
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT ELIZABETH C. KEMMERER						
Weshington, D.C. 20231						

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INTERNATIONAL SEARCH REPORT

enternational application No. PCT/US94/14426

C (Continua	tion). DOCUMENTS CONSIDERED TO BE RELEVANT		
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Y	EMBO Journal, Volume 8, Number 5, issued 1989, N. J. G. Webster et al, "The transcriptional activation function located in the hormone-binding domain of the human oestrogen receptor is not encoded in a single exon", pages 1441-1446, see entire document.		1-19
Y	E. Folkers et al, "The Retinoic Acid Receptor- β 2 Cont Separate Cell-Specific Transactivation Domains, at the	inus and in the Ligand-Binding Domain", pages 616-627, see	
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Y	Cell, Volume 68, issued 24 January 1992, R. A. Heym "9-Cis Retinoic Acid Is a High Affinity Ligand for the Receptor", pages 397-406, see entire document.	an et al, Retinoid X	1-19
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